AMENDMENTS TO THE CLAIMS/LISTING OF CLAIMS:

This listing of the claims will replace all prior versions and listings that have been previously submitted within this application:

- 1. (Currently Amended) A composition suitable for medical and surgical applications, comprising:
- a biologically compatible scaffold material having at least one irregular surface and including one of small intestine submucosa or poly(L-lactic-co-glycolic acid) (PLGA), and
- a biologically compatible light-activated adhesive, the light-activated adhesive including bovine serum albumin and a light absorber selected from at least one of red food coloring, blue food coloring and green food coloring, the light-activated adhesive also being coupled to the scaffold to form a composite, such that when the irregular surface of the composite is applied to biological tissue and the composite is activated by light energy to repair the biological tissue, the composite has a tensile strength of at least about 130% of the tensile strength of the adhesive alone.
- 2. (Original) The composition of claim 1, wherein the time-to-failure of the biological tissue repair is at least about 150% of the time-to-failure of a composite when a smooth surface of the scaffold is applied.
 - 3. (Cancelled)
 - 4. (Cancelled)
 - 5. (Cancelled)
- 6. (Currently Amended) A composition adaptable to repair biological tissue, comprising:
- a biologically compatible scaffold material, the scaffold material being selected from at least one of poly(glycolic acid), poly(L-lactic-co-glycolic acid), poly (epsilon-caprolactone), poly(ethylene glycol), poly(ortho ester)s, poly (anhydride)s, small intestine submucosa, polymerized collagen, and polymerized elastin,
 - a biologically compatible adhesive including bovine serum albumin, and
- a light absorber including one of food colorings at least one of red food coloring, blue food coloring and green food coloring, pH-indicators, water and hemoglobin, the light

absorber having a concentration of about $200 - 1000 \,\mu\text{L} / 13 \,\text{mL}$ of deionized water.

- 7. (Cancelled)
- 8. (Original) The composition of claim 6, wherein the light absorber is selected to provide a solder/interface temperature of 66 ± 3 °C.
 - 9. (Cancelled)
- 10. (Previously Presented) The composition of claim 6, wherein the light absorber concentration is about 600 μ L / 13 mL deionized water.
 - 11. (Cancelled)
 - 12. (Cancelled)
- 13. (Original) The composition of claim 6, wherein the green food coloring includes blue #1 and yellow #5.
 - 14-31. (Cancelled)
- 32. (Previously Presented) The composition of claim 1, wherein the biologically compatible scaffold material has a defined length and includes a plurality of surface irregularities spaced apart along the length of the scaffold material.
- 33. (Previously Presented) The composition of claim 32, wherein the plurality of surface irregularities are introduced to the scaffold material by at least one of molding, drilling and punching.
- 34. (Previously Presented) The composition of claim 1, wherein the scaffold material is adapted to deliver a biologically active material to a wound when the composite is applied to the wound.
- 35. (Previously Presented) The composition of claim 1, further comprising a biologically active material selected from at least one of antibiotics, anesthetics, anti-

inflammatories, bacteriostatics, bacteriocidals, chemotherapeutic agents, vitamins, antineovascular growth factors, pro-neovascular growth factors, tissue cell growth factors, hemostatic agents and thrombogenic agents.

- 36. (Previously Presented) The composition of claim 1, wherein the scaffold material provides reinforcement for wound repair in combination with the light-activated adhesive without any sutures, stapes, clips or other closure devices.
- 37. (Previously Presented) The composition of claim 1, wherein the scaffold material is adapted to provide a continuous alignment of opposing portions of biological tissue needing repair.
- 38. (Currently Amended) The composition of claim 5 1, wherein the poly(L-lactic-co-glycolic acid) has an 85:15 lactic:glycolic copolymer ratio.
- 39. (Previously Presented) The composition of 38, wherein the biologically compatible adhesive includes 50% w/v bovine serum albumin.
- 40. (Previously Presented) The composition of claim 6, wherein the biologically compatible scaffold material comprises poly(L-lactic-co-glycolic acid) having an 85:15 lactic:glycolic copolymer ratio.
- 41. (Previously Presented) The composition of claim 40, wherein the biologically compatible adhesive includes 50% w/v bovine serum albumin.
- 42. (Previously Presented) The composition of claim 6, wherein the biologically compatible scaffold material has a defined length and includes a plurality of surface irregularities spaced apart along the length of the scaffold material.
- 43. (Previously Presented) The composition of claim 42, wherein the plurality of surface irregularities are introduced to the scaffold material by at least one of molding, drilling, and punching.
 - 44. (Previously Presented) The composition of claim 6, wherein the scaffold material

is adapted to deliver a biologically active material to a wound when the composite is applied to the wound.

- 45. (Previously Presented) The composition of claim 6, further comprising a biologically active material selected from at least one of antibiotics, anesthetics, anti-inflammatories, bacteriostatics, bacteriocidals, chemotherapeutic agents, vitamins, anti-neovascular growth factors, pro-neovascular growth factors, tissue cell growth factors, hemostatic agents and thrombogenic agents.
- 46. (Previously Presented) The composition of claim 6, wherein the scaffold material provides reinforcement for wound repair in combination with the adhesive without any sutures, stapes, clips or other closure devices.
- 47. (Previously Presented) The composition of claim 6, wherein the scaffold material is adapted to provide a continuous alignment of opposing portions of biological tissue needing repair.
- 48. (Currently Amended) A composition adaptable to repair biological tissue, comprising:
- a biologically compatible scaffold material, the scaffold material being selected from at least one of poly(glycolic acid), poly(L-lactic-co-glycolic acid), poly (epsilon-caprolactone), poly(ethylene glycol), poly(ortho ester)s, poly(anhydride)s, small intestine submucosa, polymerized collagen, and polymerized elastin,
- a biologically compatible light-activated adhesive including bovine serum albumin, the light-activated adhesive including a light absorber selected from at least one of red food coloring, blue food coloring and green food coloring, the light absorber being selected to provide a solder/interface temperature of 66 + 3°C and having a concentration of about $200 1000 \,\mu\text{L} / 13 \,\text{mL}$ of deionized water.
- 49. (Previously Presented) The composition of claim 48, wherein the light absorber concentration is about 600 μ L / 13 mL deionized water.
- 50. (Previously Presented) The composition of claim 48, further comprising a biologically active material selected from at least one of antibiotics, anesthetics, anti-

inflammatories, bacteriostatics, bacteriocidals, chemotherapeutic agents, vitamins, antineovascular growth factors, pro-neovascular growth factors, tissue cell growth factors, hemostatic agents and thrombogenic agents.

- 51. (Previously Presented) The composition of claim 48, wherein the biologically compatible scaffold material comprises poly (L-lactic-co-glycolic acid) having an 85:15 lactic:glycolic copolymer ratio.
- 52. (Previously Presented) The composition of claim 48, wherein the biologically compatible adhesive includes 50% w/v bovine serum albumin.